

## Ozone production in boreal fire smoke plumes using observations

## from the Tropospheric Emission Spectrometer and the Ozone

## 4 Monitoring Instrument

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- 9 [1] We examine the photochemical processes governing the production of ozone in smoke from large Siberian fires that formed in July 2006 using colocated O<sub>3</sub> and
- 11 CO profiles as measured by the Tropospheric Emission Spectrometer as well as NO<sub>2</sub> and
- aerosol optical depths as measured by the Ozone Monitoring Instrument. The Real-Time
- Air Quality Model (RAOMS) is used to explain the observed variations of O<sub>3</sub>. Enhanced
- levels of ozone up to 90 parts per billion (ppbv) are observed near and away from
- the Siberian fires  $(60^{\circ}\text{N and }100^{\circ}\text{E})$  when sunlight and  $NO_x$  are available. We also observe
- significantly low O<sub>3</sub> amounts (less then 30 ppbv) in the smoke plume from Siberian fires
- in conjunction with optically thick aerosols. Despite this wide variance in observed ozone
- values, the mean ozone value for all observations of the smoke plume is close to
- background levels of approximately 55 ppbv in the free troposphere. Using RAQMS we
- show that optically thick aerosols in the smoke plume can substantially reduce the
- 21 photochemical production of ozone and this can explain why the observed mean ozone
- 22 amount for all plume observations is not much larger than background values of 55 ppbv.
- However, the anonymously low ozone amounts of 30 ppbv or less point toward other
- unresolved processes that reduce ozone below background levels in the plume.
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### 1. Introduction

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[2] Long-range transport of smoke emissions from boreal fires can increase the atmospheric abundance of pollution over population centers. Prior studies [Wotawa and Trainer, 2000; Forster et al., 2001] have shown, for example, that transport of emissions from the Canadian boreal fires can significantly increase atmospheric abundances of carbon monoxide (CO), aerosols, and ozone (O<sub>3</sub>) over North American and European population centers. Bertschi and Jaffe [2005] showed, using satellite observations of aerosols and global aerosol transport model, that Siberian fire emissions were the primary source of three air pollution events off the coast of Washington State in 2003.

[4] Extensive fires burned in Siberia during the summer 62 of 2006 as shown in Figure 1 which shows image from the 63 Moderate Resolution Imaging Spectroradiometer (MODIS). 64 As observed by both the Tropospheric Emission Spectrom-65 eter (TES) and the Ozone Monitoring Instrument (OMI), 66

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<sup>[3]</sup> Boreal fires play an important role in the magnitude 41 and interannual variability of tropospheric CO in the North- 42 ern Hemisphere [e.g., Novelli et al., 2003; Edwards et al., 43 2004; Kasischke et al., 2005; Pfister et al., 2005; Nedelec et 44 al., 2005]. Several studies indicate that boreal fires can 45 impact summertime O<sub>3</sub> over northwestern North America 46 [Jaffe et al., 2004; Morris et al., 2006], the central North 47 Atlantic [Lapina et al., 2006] and Europe [Simmonds et al., 48 2005]. The understanding of how boreal fires impact 49 tropospheric O<sub>3</sub> is difficult, however because O<sub>3</sub> production 50 in boreal fires is highly variable [Mauzerall et al., 1996; 51 Lapina et al., 2006; Val Martin et al., 2006; Real et al., 52 2007]. Real et al. [2007] reported significant O<sub>3</sub> variations 53 in fire plumes depending upon the photochemical history of 54 each plume. Val Martin et al. [2006] studied the impact of 55 boreal fires in northern North America on the levels of black 56 carbon (BC) aerosols, nitrogen oxides (NO<sub>x</sub>) and O<sub>3</sub> down- 57 wind from North America. They noted that the boreal 58 wildfire emissions significantly contributed to the NO<sub>x</sub> 59 and O<sub>3</sub> budgets in the lower free troposphere over the 60 central North Atlantic during the summer of 2004.

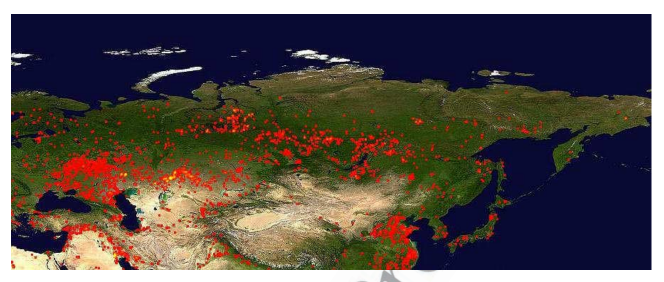
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**Figure 1.** Spatial distribution of the Siberian fires detected by MODIS during the period 20–26 July 2006.

the plumes from these fires stretched across Siberia and the Pacific Ocean. The availability of satellite observations of CO, O<sub>3</sub>, aerosols optical depth (AOD), and nitrogen dioxide (NO<sub>2</sub>), for this period provides a valuable opportunity to evaluate our understanding of factors controlling boreal fire emissions, their impact on atmospheric chemistry, and the transport of the ozone produced in this plume globally. This study examines O<sub>3</sub> produced and transported during July 2006 Siberian fire smoke plumes as observed from space-based observing platforms such as the Tropospheric Emission Spectrometer (TES) and the Ozone Monitoring Instrument (OMI) on the EOS-Aura satellite [Schoeberl et al., 2006].

## 2. Satellite Observations Used in Analysis

[5] The TES instrument is an infrared Fourier transform spectrometer that measures the thermal emission of the Earth's surface and atmosphere over the spectral range  $650-2250~\rm cm^{-1}$ . It was designed to provide simultaneous vertical profile retrievals of tropospheric  $O_3$ , CO and other trace gases on a global basis [Beer et al., 2001; Beer, 2006]. The nadir footprint is 5.3 km across the spacecraft ground track and 8.5 km along track for the 16-detector average [Beer et al., 2001]. TES has two basic science operating modes: Global Survey and Special Observations. Global Surveys are conducted every other day while special observations are taken as needed in between Global Surveys. We used global survey observations of TES  $O_3$  and CO obtained between 20 July and 12 August 2006 with a nadir sampling of  $\sim 1.6^{\circ}$  spacing along the ground track.

[6] The analysis presented here utilizes TES version 003 data [Osterman et al., 2005]. An overview of the TES retrieval algorithm and error estimation are discussed by Bowman et al. [2006] and the characterization of errors and vertical information for individual TES profiles are discussed by Worden et al. [2004] and Kulawik et al. [2006]. The vertical resolution of TES nadir O<sub>3</sub> retrievals is about 6 km for cloud-free scenes, with sensitivity to both the

lower and upper troposphere [Worden et al., 2004; Bowman 105 et al., 2006]. To date, TES tropospheric O<sub>3</sub> validation has 106 been conducted through comparisons with ozonesondes 107 [Worden et al., 2007] and lidar [Richards et al., 2008]. 108 These validation studies show that TES O<sub>3</sub> estimates are 109 typically biased high in the upper troposphere by approx- 110 imately 10%. Nassar et al. [2007] shows that TES O<sub>3</sub> is 111 biased high by 3–10 ppb in the upper troposphere.

#### **2.2. OMI** 11

[7] OMI is an ultraviolet and visible nadir solar back- 114 scatter imaging spectrograph which provides nearly global 115 coverage in one day with a nadir spatial resolution of  $13 \times 116$   $24 \text{ km}^2$ . OMI measures solar irradiance and Earth radiances 117 in the wavelength range of 270 to 500 nm with a spectral 118 resolution of about 0.5 nm. These radiances are used for 119 estimating tropospheric column amounts of  $NO_2$ , HCHO, 120  $SO_2$ , AOD, as well as total  $O_3$  amount [Levelt et al., 2006]. 121

[8] The analysis in this study utilizes OMI aerosol optical 122 depth and NO<sub>2</sub> Level 2 version 3 data products. The OMI 123 Level 2 geolocated geophysical parameters (O<sub>3</sub>, NO<sub>2</sub>, SO<sub>2</sub>, 124 BrO, HCHO, OClO, Aerosol, and Cloud) data products are 125 at full instrument resolution, one orbit per file (http:// 126 disc.sci.gsfc.nasa.gov/data/datapool/OMI/). The basic algo-127 rithm for the retrieval of AOD and NO<sub>2</sub> from OMI data are 128 described by *Torres et al.* [2002], *Boersma et al.* [2007], and 129 *Bucsela et al.* [2006], respectively. The NO<sub>2</sub> product was 130 successfully validated with DC-8 aircraft NO<sub>2</sub> vertical 131 profiles [*Boersma et al.*, 2008].

### 3. Modeling Tool Used in Analysis: Real-Time Air 134 Quality Modeling System 135

[9] Chemical and aerosol analyses from the Real-Time Air 136 Quality modeling System (RAQMS) and ensemble wild fire 137 trajectories are used to examine the different processes 138 influencing the evolution of trace gases (e.g., O<sub>3</sub> and CO) 139 within fire plumes during the 2006 Siberian boreal fires 140 event. RAQMS is a unified (stratosphere/troposphere), 141 online (meteorological, chemical, and aerosol) modeling 142

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system which has been developed for assimilating satellite observations of atmospheric chemical composition and providing real-time predictions of trace gas and aerosol distri-145 butions [Pierce et al., 2003, 2007; Kittaka et al., 2004]. The 146chemical formulation follows a family approach with parti-147 tioning on the basis of photochemical equilibrium approx-148 imations. The nonmethane hydrocarbon (NMHC) chemical 149 scheme is based on the carbon bond lumped structure 150 approach [Pierce et al., 2007]. Photolytic rates are calculated 151 using the Fastj2 method [Bian et al., 2003]. The RAQMS 152 aerosol model incorporates online aerosol modules from 153 GOCART [Chin et al., 2002, 2003] and a sulfate-nitrate-154 ammonium thermodynamic equilibrium model [Park et al., 155 2004]. Nine aerosol species (SO<sub>4</sub>, NO<sub>3</sub>, NH<sub>4</sub>, hydrophobic 156 organic carbon (OC), hydrophilic OC, hydrophobic BC, 157 158 hydrophilic BC, dust, sea-salt) are transported. RAQMS biomass burning emissions use twice daily ecosystem/severity based emission estimates coupled with Moderate-Resolution Imaging Spectroradiometer (MODIS) Rapid Response 161 fire detections [Al-Saadi et al., 2008]. Total direct carbon 162 emissions are calculated as the product of area burned and 163 the ecosystem- and severity-specific carbon consumption 164estimates. Ecosystem-dependent carbon consumption databases for three classes of fire severity (low, medium, and high) are considered. Fire weather severity is estimated 167 using the U.S. Forest Service Haines Index, which considers 168 atmospheric moisture and thermal stability [Haines, 1988]. 169 Emissions of other species are determined by combining 170 171 published emission ratios for different ecosystems [Cofer et 172 al., 1991; Andreae and Merlet, 2001]. During the chemical and aerosol assimilation cycle the RAQMS meteorological 173 forecasts are reinitialized from NOAA Global Forecasting 174 System (GFS) analyses at 6 h intervals. 175

[10] The RAQMS chemical analysis used in the current study is from a retrospective 9-month (February-October 2006) 2 × 2° assimilation that includes assimilation of cloud cleared OMI total column O<sub>3</sub> measurements and O<sub>3</sub> and CO profiles from TES nadir measurements. MODIS onboard the Aqua satellite [Remer et al., 2005; Davies et al., 2004] AOD was assimilated during the period from 15 to 31 July to provide observational constraints for the investigation of the influences of aerosol loading on Siberian wild fire photochemistry. The MODIS AOD assimilation cycle was initialized from a Global Modeling and Assimilation Office (GMAO) aerosol forecast provided by Arlindo da Silva (NASA/GSFC). For the wild fire case studies only anthropogenic and biomass burning sources of carbonaceous aerosols were considered. Other aerosol species were passively advected. During the MODIS assimilation cycle, masses of all aerosol species were adjusted within each model layer on the basis of the total AOD analysis increment and the relative contribution of each aerosol species to the total layer extinction. The MODIS AOD compares well with OMI AOD observations also used in this study with a correlation of approximately 0.7 between the two data sets but with OMI AOD showing slightly higher values than MODIS of about 0.2 AOD [Ahn et al., 2008].

#### 200 4. Methodology

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[11] During mid-July 2006, multiple forest fires were recorded across the central Siberian Plateau between the north-flowing Lena River and Lake Baikal along 95°E 203 longitude and 60°N latitude (http://earthobservatory.nasa. 204 gov/NaturalHazards/natural\_hazards\_v2.php3?img\_id= 205 13728). The Spatial distribution of the Siberian fires 206 detected by MODIS during the period from 20 to 26 July 207 2006 is shown in Figure 1. The substantial fire hot spots 208 located over Siberia from the MODIS fire counts data 209 between 55 and 70°N are also shown in Figure 2 (marked 210 as red) on 24 July 2006, which was the day of the peak 211 Siberian wildfire emissions, as shown by MODIS fire 212 counts [Davies et al., 2004].

[12] In order to examine the production of O<sub>3</sub> within the 214 smoke plumes from these fires we first needed to identify 215 the plume signature. TES and OMI observations from mid- 216 July to mid-August 2006 were used. TES measurements 217 with CO abundances greater than 120 ppb [e.g., Wofsy et al., 218 1992; Mauzerall et al., 1996] were identified as potentially 219 impacted by upwind boreal fires. Colocated AOD from 220 OMI were used as corroborating evidence. Images from 221 MODIS were acquired during this period to compare with 222 the satellite outputs as further corroborating evidence. We 223 used backward trajectories from these observations of 224 enhanced CO and aerosols to the fire source as well as 225 forward trajectories from the fire source using both the 226 FLEXPART model [Stohl et al., 1998] and RAQMS to 227 ensure that the observed air parcels came from the Siberian 228 fires of interest.

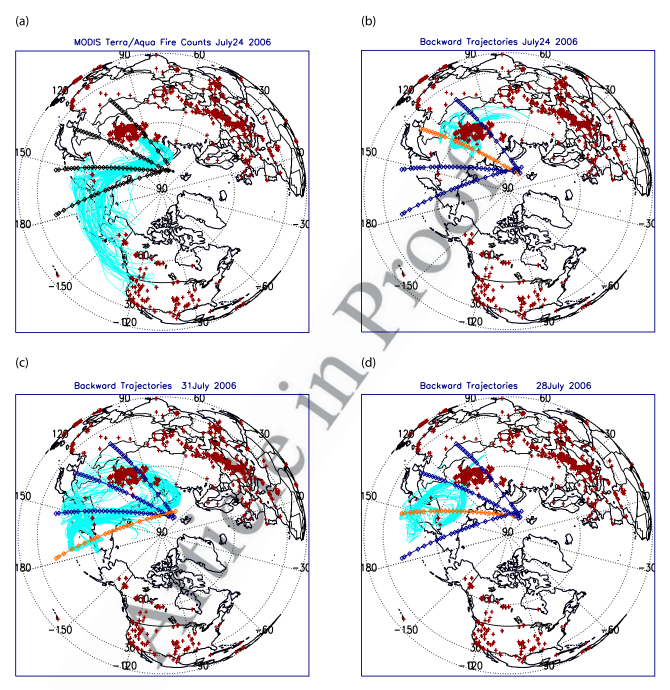
#### Discussions

#### 5.1. Siberian Boreal Fire Emissions

[13] The MODIS data revealed intense fire activity in the 232 Siberian region during July 2006, near 95°E and 65°N as 233 shown in Figures 1 and 2 (marked as red). Figure 2a shows 234 the FLEXPART 10-day forward trajectories from this fire 235 starting on 24 July 2006. The light blue lines represent the 236 forward trajectories for a plume starting from a TES orbit 237 around 98°E and 62°N with altitude levels between 0 and 238 15 km. Figures 2b, 2c, and 2d shows the FLEXPART 5-day 239 backward trajectories (light blue lines) starting from a TES 240 track location (shown as orange), corresponding to Figures 3, 241 4, and 6, respectively. The back trajectories in Figures 2b, 242 2c, and 2d suggests that air sampled on the selected TES 243 orbits is mainly from the Siberian fires as the trajectories 244 show significant recirculation in and around the fire source. 245 The forward trajectories in Figure 2a indicate large plumes 246 of smoke transported from northeastern Siberia to across the 247 Pacific during late July, providing opportunities for many observations of the plume by TES and OMI.

#### 5.2. TES and OMI Observations for Siberian Smoke Plume

[14] Because the smoke from the fires of interest travels 252 eastward and the Aura satellite travels along a polar orbit 253 there are many opportunities to observe cross section of the 254 fire plumes. We first show four examples showing vertical 255 cross sections of CO and O3 values across fire plumes. The 256 average CO and O<sub>3</sub> mixing ratios is also presented along 257 these vertical cross sections. As discussed earlier, in order to 258 ensure that the observed air parcels are only related to 259 smoke from the fires, the CO values from each observation 260 are averaged together over the pressure (>400 hPa) range 261



**Figure 2.** (a) The 10-day forward trajectories (blue lines) from peak Siberian wildfire emissions for a plume starting around 98°E and 62°N on 24 July 2006. (b-d) Five-day backward (blue lines) trajectories starting from a TES track (shown as orange), corresponds to Figures 3, 4, and 6, respectively. The trajectories represent the five different sets of latitude and longitudes pairs over the TES overpass and altitude levels between 0 and 15 km. Locations shown in red (marked as crosses) indicate the MODIS fire retrievals on 24 July 2006. The orbit tracks correspond to TES measurement locations chosen as examples in this study, as shown in Figures 4b, 6b, 3b, and 5b, respectively, starting from west to east.

for which CO values are larger than 120 parts per billion by volume (ppbv). The ozone values are also averaged using similar criteria.

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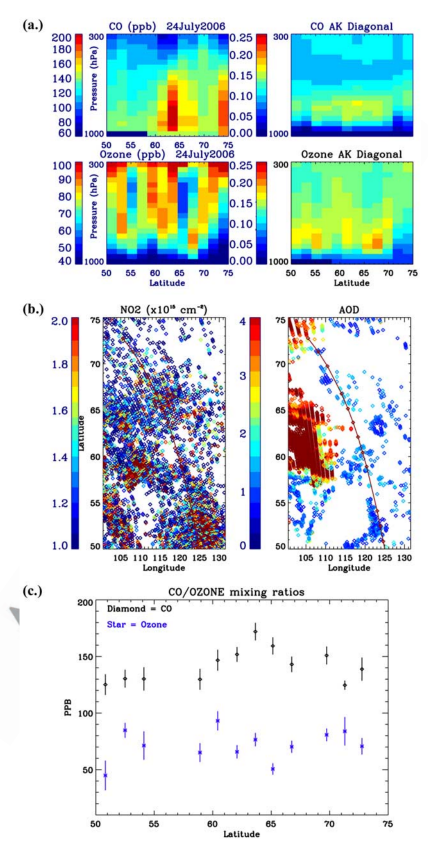
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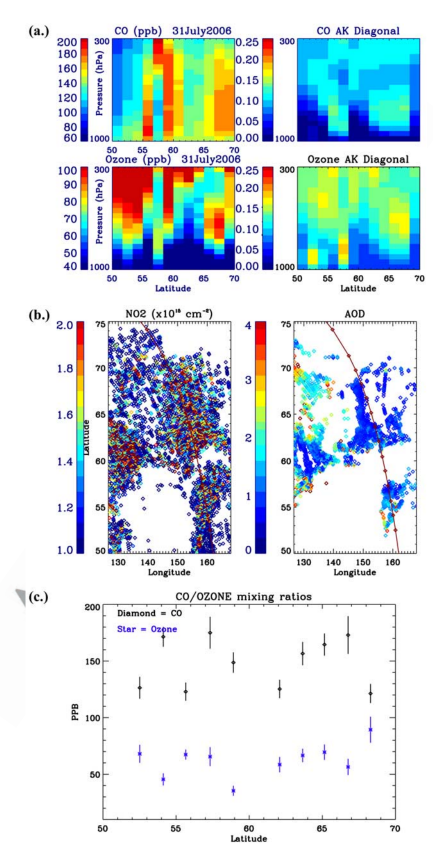
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[15] Figure 3a shows CO and  $O_3$  profiles from TES on 24 July 2007 near the Siberian fire source (100–130°E). The location of the Siberian fires are observed by MODIS to be approximately between  $90-110^{\circ}E$  and  $60-70^{\circ}N$  (Figures 1

and 2). The vertical profiles of CO in Figure 3a shows 269 values ranging from background levels (~80 ppb) to 270 enhanced levels (between 120 and 250 ppb). Back trajec-271 tories from the regions of high CO (Figure 2b) indicate that 272 observed air parcels have recirculated in and around the fire 273 and therefore indicate the high CO is related to the fire 274 emissions.



**Figure 3.** (a) TES CO,  $O_3$ , and Averaging Kernel (AK) diagonals, (b) AOD and  $NO_2$  tropospheric column amounts as observed from OMI, and (c) latitudinal variations of CO and  $O_3$  mixing ratios averaged for CO >120 ppb and pressure >400 hPa for the plume values near the Siberian fires at 100–130°E on 24 July 2006. Overlaying the AOD and  $NO_2$  (Figure 3b) is the TES orbit track (red curve) with the locations of the TES observations indicated by diamonds.



**Figure 4.** Same as in Figure 3 but for the plume away from the Siberian fires at 135–170°E on 31 July 2006.

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[16] The vertical resolution of the TES ozone and CO estimates is approximately 6 km [Worden et al., 2006]. However, it is appropriate to show these vertical profiles because the plume height depends on altitude and the sensitivity of the estimates depend on altitude. The sensitivity of the TES estimates are indicated by the diagonal of the averaging kernel matrices shown to the right of the CO and O<sub>3</sub> estimates. The diagonal of the averaging kernel matrix is the sensitivity of the O<sub>3</sub> or CO estimate at the indicated pressure level to the actual amount of the species at the same pressure level. The greatest sensitivity of the CO and O<sub>3</sub> estimates is in the free troposphere between 400 hPa and 800 hPa. Peak values of the O<sub>3</sub> and CO averaging kernel diagonals correspond to peak values of O<sub>3</sub> and CO, and the largest variations of O3 relative to background values of approximately 55 ppbv in the free troposphere. Consequently it is reasonable to assume that observed variations in O<sub>3</sub> and CO amounts in the free troposphere are at similar altitudes.

[17] Figure 3b shows aerosol optical depths as observed by OMI between 100 and 130°E longitude and 50 and 75°N latitude. The TES orbit track is also shown as a red curve over AOD values and the location of the TES observations are shown as diamonds overlaying the orbit track. The aerosols provide corroborating evidence that the observed air parcel interacted with the boreal fire plume. In fact, the largest values of AOD between 100 and 110°E and 60-70°N correspond to the fire locations as seen by the MODIS fire count data in Figure 1. Back trajectories from the locations of the TES observations (Figure 2b) indicate that the air parcels observed by TES had intersected the location of the fire approximately three days earlier. Enhanced O3 is observed anywhere from 50°N to 74°N. NO<sub>2</sub> values of approximately  $2 \times 10^{15}$  molec/cm<sup>2</sup> and relatively low AOD (AOD < 0.5) as seen in Figure 3b, indicate the availability of O<sub>3</sub> precursors and sunlight for photochemical production of O<sub>3</sub>.

[18] In order to better examine O<sub>3</sub> and CO variations within the plume we next average the CO values over those altitudes where CO is larger than 120 ppbv and the diagonal of the averaging kernel is larger than 0.05 and the sum of the diagonals (also known as the degrees of freedom for signal) is larger than 0.5 for CO in the troposphere. The O<sub>3</sub> concentrations averaged over these same altitudes are shown in Figure 3c for observations that meet the above criteria. The total error estimates, calculated using the total error for each profile, averaged over the selected altitudes [Worden et al., 2006], is also shown. We find the total error for these averaged quantities are about 8–12% for O<sub>3</sub> and 6-10% for CO; these uncertainties are sufficient for resolving the observed O<sub>3</sub> and CO variations. For this plume cross section, we observe values ranging from 130 to 180 ppbv for CO and approximately 35 to 90 ppbv for O<sub>3</sub>. There appears to be no correlation between the O<sub>3</sub> and CO amounts.

[19] These same quantities are shown in Figure 4a for a region further away from the fire (approximately  $50^{\circ}$  in Longitude) and at a later date (31 July 2007). Enhanced levels of CO and AOD (AOD > 0.1) as well as back trajectories (Figure 2c) indicate the enhanced CO levels are due to the Siberian fire emissions. As seen in Figure 4c, enhanced  $O_3$  of up to 70 ppbv is observed in regions where CO is moderately enhanced (between 120 and 200 ppb) and AOD is moderately enhanced (AOD  $\sim$  0.5) along with the

availability of NO<sub>2</sub>. However, there are also a couple of 338 observations with low ozone values which likely indicate 339 that different parts of this large plume, covering up to 15° in 340 latitude, have different chemical histories. We explore these 341 low ozone values next.

Figures 5a and 6a show cross sections of the fire plume 344 where  $O_3$  is low relative to background values of approx-345 imately 55 ppb in the free troposphere. Figure 5a and 5b 346 shows CO,  $O_3$ , AOD and  $NO_2$ , respectively directly over 347 the region where the fires are burning on 24 July 2006. CO 348 is observed with values exceeding 300 ppb (with the color 349 scale on the top left panel saturating at 200 ppb) and the 350 aerosol optical depth exceeding 4.  $NO_2$  is observed with 351 values ranging between  $1 \times 10^{15}$  and  $2 \times 10^{15}$  molec/cm<sup>2</sup>. 352 Ozone values ranging from 30 to 65 ppbv are observed. The 353 lower ozone values occur in regions where AOD is optically 354 thick (>3). Note that the high AOD values do not affect the 355 TES estimates of CO and  $O_3$  because thermal infrared 356 radiation is not absorbed much by aerosols produced in fire 357 [Kirchstetter et al., 2004].

[21] This behavior of low  $O_3$  within a plume with 359 significant aerosol optical depth and significantly enhanced 360 CO amounts (greater the 200 ppb) is also observed (Figure 361 6a) four days later (on 28 July 2008) for a plume that is  $40^\circ$  362 away from the fire source at approximately  $100^\circ$ E. In both, 363 the 24 and 28 July cases there is significant variation of  $O_3$  364 across the plume, with low  $O_3$  usually occurring in plume 365 cross sections with high AOD (AOD > 3) (Figure 6b). 366 Because this behavior is observed both over and away from 367 the fire, where temperature conditions are expected to be 368 different, we do not believe that incorrect estimates of 369 temperature will affect these  $O_3$  retrieval results.

[22] To examine these relationships further we compute 371 these same values as shown in Figures 4c through 8c for 372 approximately 30 plume cross sections of mid-July-mid- 373 August 2006 where the air sampled originated from the 374 Siberian smoke plumes. The mean O<sub>3</sub> and CO is computed 375 for each cross section along with the root mean square of 376 the O<sub>3</sub> variability across the plume. These values are shown 377 for these cross sections in Figure 7. The symbols in Figure 7 378 are the mean values of CO and O<sub>3</sub> in the plume cross 379 section and the bars show the RMS of the O<sub>3</sub> in the cross 380 section. We observe that O<sub>3</sub> can vary significantly, with 381 values ranging from 20 ppbv to 90 ppbv for a wide range of 382 CO values. Despite this large variance, the mean O<sub>3</sub> for all 383 observations is about the same as the background value of 384 about 55 ppbv observed at similar latitudes but without 385 enhanced CO. This suggests that while enhanced O<sub>3</sub> abun- 386 dances are produced in this boreal fire plume, the average 387 net O<sub>3</sub> production in the plume is small which is consistent 388 with prior observations of aged O<sub>3</sub> plumes using tower and 389 aircraft observations [Real et al., 2007; Mauzerall et al., 390 1996; Val Martin et al., 2006]. 391

#### 5.3. RAQMS Analysis

[23] The chemical evolution of the Siberian boreal fires is 393 further explored by sampling the RAQMS chemical analysis along the ensemble fire trajectories to understand the 395 different processes and time evolution of trace gases within 396 the plumes, with a particular focus on the evolution of  $O_3$  397 and  $O_4$  and  $O_4$  are the 398 are trajectories to understand the 398 are trajectories to under

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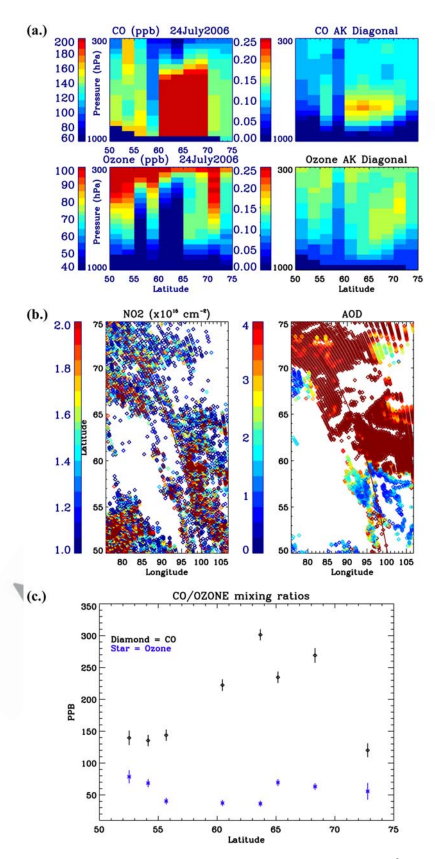
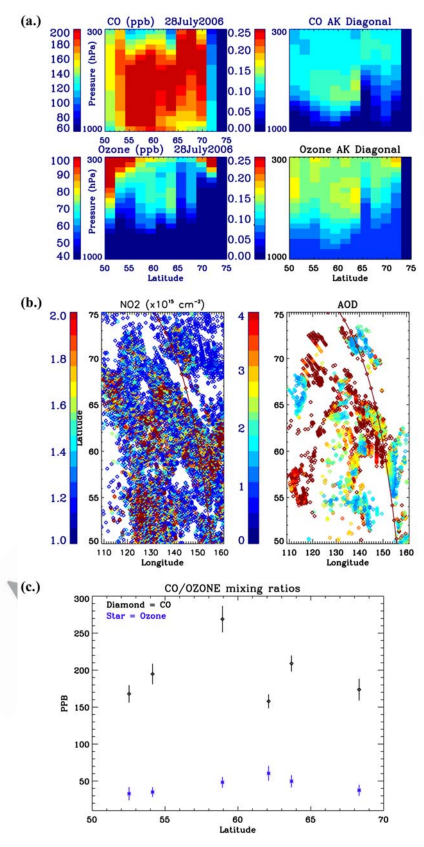


Figure 5. Same as in Figure 3 but for the plume near the Siberian fires at 80–110°E on 24 July 2006.



**Figure 6.** Same as in Figure 3 but for the plume away from the Siberian fires at 130–160°E on 28 July 2006.

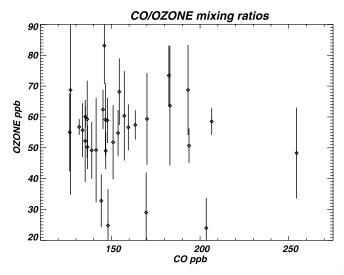


Figure 7. Scatterplot for CO and O<sub>3</sub> mixing ratios for all plume observations averaged for the Siberian fire period (mid-July-mid-August 2006).

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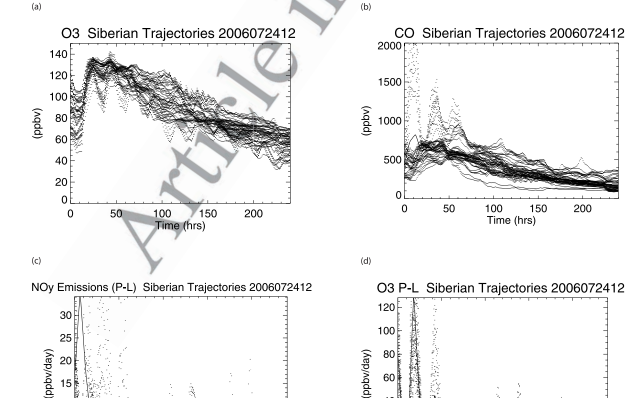
Time (hrs)

impact of aerosols on O<sub>3</sub> production rates within the wild 399 fire plumes we conducted two simulations, one with and one 400 without BC and OC aerosols in the photolysis calculations. 401

### 5.3.1. RAQMS Chemistry Run Without Aerosols

[24] Figure 8 represent results from a photochemical 403 calculation of O<sub>3</sub> in the RAQMS model in which an 404 ensemble of 10-day forward trajectories samples the model 405 output starting on 24 July 2006 for Siberia at 100°E, 60°N 406 location. This run was conducted for the chemistry only 407 simulation from RAQMS, that is, without aerosols in the 408 photolysis calculations. On the basis of the RAQMS wild 409 fire emission estimates, this time period is associated with 410 the highest wild fire emission rates for this Siberian fire. 411 The time evolution of  $O_3$ , CO concentrations,  $NO_v$  (or  $NO_x$  412 per day) and net O<sub>3</sub> production rates (production-loss) is 413 shown in Figure 8 along the smoke plume from this 10 day 414 forward trajectory run. The time histories for each ensemble 415 member (dots) and ensemble mean (solid line) are shown. 416 There is a large increase in O<sub>3</sub> concentrations in the fresh 417 part of plume (ensemble mean goes from 80 to 130 ppb 418 within the first 24 h). This increase in O<sub>3</sub> concentrations 419 is due to large daytime net production of O<sub>3</sub> (exceeding 420 120 ppb/day 24 hrs after the start of the trajectory calcula- 421 tion) associated with high NO<sub>2</sub> mixing ratios (net NO<sub>2</sub> 422

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**Figure 8.** The 10-day forward trajectory simulation without aerosols initialized with 24 July 2006 upwind data and daily fire emissions (Tg/day) from real time MODIS fire counts at Siberia showing time evolution of (a) O<sub>3</sub>, (b) CO concentrations, (c) NO<sub>v</sub>, and (d) O<sub>3</sub> production rates.

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Time (hrs)

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(a) (b)

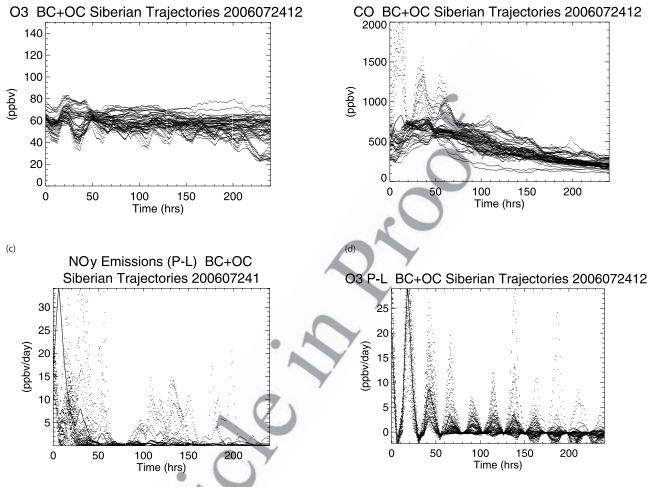


Figure 9. Same as in Figure 8 but with aerosols.

emissions reach 34 ppb/day). The peak O<sub>3</sub> mixing ratios within the fresh wild fire plume in the model of nearly 140 ppb is significantly larger than observed by TES which shows peak values of 90 ppb in the vicinity of the wild fire plume (see Figure 3a).

# 5.3.2. RAQMS Sensitivity Run: Aerosols Impact on Ozone Production (High Aerosol Loading)

[25] A second ensemble of 10-day forward trajectories starting on 24 July 2006 Siberia is shown in Figure 9 for a RAQMS chemistry simulation that also includes BC and OC aerosols in the photolysis calculations. In contrast to the chemistry only simulation, the RAQMS model including aerosols shows only slight increases in O<sub>3</sub> concentrations in the fresh part of the plume with the ensemble mean ranging from 60 to 70 ppb within the first 24 hrs. This slight increase in O<sub>3</sub> concentrations is due to enhanced daytime net O<sub>3</sub> production (which was about 29 ppb/day at 24 hrs). This represents a fourfold decrease in net O<sub>3</sub> production relative to the model simulation without aerosols and suggests that optically thick aerosols (AOD > 3) during the most intense phase of the Siberian wild fire event significantly inhibit photolysis and hence greatly modifies the O<sub>3</sub> production; this conclusion is consistent with the results of Real et al. [2007] who also discussed the effects

of aerosols on reduced  $O_3$  production. However, neither 447 simulation is able to replicate the low  $O_3$  observations of 448 about 30–40 ppb observed by TES, which may be due to 449 other chemical processes at the fire source such as aerosol 450 surface chemistry [e.g., *Val Martin et al.*, 2006; *Real et al.*, 451 2007] or  $O_3$  destruction due to the titration of  $NO_x$  within 452 the fresh plume [*Crutzen and Bruhl*, 2001].

#### 6. Discussion and Conclusions

[26] Ozone production in the July 2006 Siberian boreal 456 fire is examined using synchronous tropospheric observations of  $O_3$  and CO from TES and observations of aerosol 458 optical depth and  $O_2$  column abundances from OMI. 459 These observations show that Siberian biomass burning 460 emissions can produce elevated  $O_3$  within the fire plume. 461 However,  $O_3$  abundances in the Siberian boreal forest fire 462 plumes are highly variable, with some plumes showing  $O_3$  463 enhancements of up to 90 ppb and others showing no 464 enhancement or even  $O_3$  depletion, with abundances of 465 30 ppb, much lower than background tropospheric values of 466 about 55 ppb.

[27] We investigated the impact of aerosols on O<sub>3</sub> pro- 468 duction rates within the wild fire plumes using the RAQMS 469

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model. In the absence of aerosols, the RAOMS model predicts up to 120 ppb of O<sub>3</sub> in the fire plume. Accounting for the presence of optically thick aerosols by assimilating in the model AOD data from the MODIS instrument reduced the photolysis rates in the model. As a result, in regions with optically thick aerosols, the model predicted a 475 significant decrease in the net production of O<sub>3</sub>, from about 476 120 ppb/day to 30 ppb/day within 24 hrs of the plume 477 emission. O<sub>3</sub> concentrations in the model simulation with 478 the assimilated AOD were approximately 60 ppb which is 479 consistent with the mean ozone for all plume observations. 480 Reduced photolysis due to aerosols, and, by the same 481 reasoning, clouds, could therefore explain the lack of 482 enhanced O<sub>3</sub> levels seen in the boreal fire plume produced 483 by the Siberian fires. However, the anomalously low O<sub>3</sub> 484 abundances of 30 ppb are also observed indicating that additional O<sub>3</sub> loss mechanisms, such as NO<sub>2</sub> titration or aerosol surface chemistry, could be important in determin-487 ing the observed O<sub>3</sub> abundances. Combining these satellite 488 observations with in situ observations of O<sub>3</sub> and its precursors such as those from the 2008 Arctic Research of the 490 Composition of the Troposphere from Aircraft and Satellites (ARCTAS: http://www.espo.nasa.gov/arctas/) aircraft campaign should greatly improve understanding of ozone production and loss mechanisms in boreal fire smoke plumes and their impact on global atmospheric composition. 495

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